

This Page Is Inserted by IFW Operations
and is not a part of the Official Record

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images may include (but are not limited to):

- BLACK BORDERS
- TEXT CUT OFF AT TOP, BOTTOM OR SIDES
- FADED TEXT
- ILLEGIBLE TEXT
- SKEWED/SLANTED IMAGES
- COLORED PHOTOS
- BLACK OR VERY BLACK AND WHITE DARK PHOTOS
- GRAY SCALE DOCUMENTS

IMAGES ARE BEST AVAILABLE COPY.

**As rescanning documents *will not* correct images,
please do not report the images to the
Image Problem Mailbox.**

Remarks/Arguments

The elected claims are directed to a method for identifying a HCV RNA-dependent RNA polymerase inhibitor using an *in vitro* assay. The present application provides data demonstrating that NS5B does in fact produce RNA-dependent RNA polymerase activity and that sufficient polymerase activity can be produced *in vitro* to identify a polymerase inhibitor. The application provides motivation, and examples of techniques, for the skilled artisan to move beyond mere identification and characterization of polymerase activity into assaying for polymerase inhibitors.

35 U.S.C 103 (Obviousness)

Claims 12, 14, 16, 17 and 18 stand rejected as allegedly obvious based on either (1) Behrens et al. (EMBO 15(1): 12-22, January 1, 1996); (2) Al et al. (Hepatology 22(4 part 2): 331A, October 1995); or (3) Tomei et al. (Journal of Virology 67(7): 4017-4026, July 1993). These rejections are respectfully traversed.

The prior office action notes that the rejections based on Behrens et al. and Al et al. may be overcome by filing a translation of the claimed priority document. Enclosed is a copy of a certified translation of the priority document. A copy of the translation was also filed in the parent application.

The office action argues that Tomei et al. suggests that NS5B may act as a viral replicase based on the presence of a GDD sequence. Tomei et al. is argued to provide motivation for further characterizing the function and roles of NS5B encoded proteins, determining whether proteolytic processing affects NS5B protein product, characterizing the specific mechanism of RNA-dependent RNA polymerase activity, and being used to identify potential therapeutics.

It is respectfully submitted that Tomei et al., fails to provide a reasonable expectation of success that NS5B can be used in an *in vitro* assay for identifying a HCV RNA-dependent RNA polymerase inhibitor. Tomei et al. speculates that NS5B **may** act as a viral replicase and points out that the HCV NS5 region is processed differently than flavivirus NS5B:

The NS5 region of the HCV polyprotein is cleaved into two smaller products of 47 and 65 kDa; **the processing of this region therefore differs from that of flavivirus NS5**, which is released from the polyprotein precursor as a single protein of 110 kDa. The GDD consensus sequence characteristic of RNA-

dependent RNA polymerases is located in NS5b (residues 2736 to 2738), indicating that this protein **may** act as a viral replicase during HCV-specific RNA synthesis (17). However, NS5a could also have a function in the replication of the viral genome, acting as a component of the replication complex involved in the reaction. [Emphasis added.]

Tomei et al., fails to provide any data measuring RNA-dependent RNA polymerase activity from NS5B *in vitro*.

Absent data demonstrating that NS5B provides RNA-dependent RNA polymerase activity, the skilled artisan would have no motivation to perform an assay measuring RNA-dependent RNA polymerase activity from NS5B *in vitro* to assay for a HCV polymerase inhibitor. Not only is there uncertainty as to (1) whether NS5B provides RNA-dependent RNA polymerase, but there is also uncertainty as to (2) whether NS5B, if it encodes the polymerase, can be used in an *in vitro* assay to generate sufficient activity to identify inhibitors.

Double Patenting

The pending claims stand rejected for double patenting based on U.S. Patent No. 6,383,768 ('768 patent). Claims 12, 14, 16, 17 and 18 stand rejected under obviousness type double patenting. Claims 13, 15 and 19 stand rejected for same invention double patenting. The rejections are respectfully traversed.

With respect to obviousness-type double patenting, as noted by the Examiner, applicants may overcome the rejection by filing a terminal disclaimer. Because of the possibility that the claims may be amended in the future, applicants have not yet filed a terminal disclaimer.

Pending claims 13, 15 and 19 were rejected for same invention double patenting based on claims 8, 15 and 11 of the '768 patent. The differences between these claims can be illustrated by the difference between pending claim 13 and '768 patent claim 8. Pending claims 15 and 19 each depend on claim 13. '768 patent claims 15 and 11 depend from '768 patent claim 8 and provide similar limitations as pending claim 15 and 19.

Pending claim 13 differs from the '768 patent claim 8, for example, in that pending claim 13 indicates that NS5B is the only HCV protein present during said incubation. In contrast, the '768 patent claim 8 characterizes the source of NS5B provided to an assay composition as "said

NS5B is provided to said composition from a preparation wherein said NS5B is the only HCV protein present . . .”.

Accordingly the claims are in condition for allowance. Please charge deposit account 13-2755 for fees due in connection with this amendment. If any time extensions are needed for the timely filing of the present amendment, Applicants petition for such extensions and authorize the charging of deposit account 13-2755 for the appropriate fees.

Respectfully submitted,

By Sheldon Heber
Sheldon O. Heber
Reg. No. 38,179
Attorney for Applicants

Merck & Co., Inc.
P.O. Box 2000
Rahway, NJ 07065-0907
(732) 594-1958

TCNY

TRANSLATION COMPANY
OF NEW YORK, Inc.

8 S. Maple Avenue, Marlton, NJ 08053 (856) 983-4733



CERTIFICATE OF ACCURACY

STATE OF NEW JERSEY)
COUNTY OF BURLINGTON) SS.:

Elsbeth LUSSI, being duly sworn, deposes and says that she is the President of Translation Company of New York, Inc. and that she knows Susan Geddes, who is thoroughly familiar with the Italian and the English language and that (s)he translated the attached document relating to:

Patent Application - "Method for Reproducing in vitro the RNA-Dependent RNA Polymerase and Terminal Nucleotidyl Transferase Activities Encoded by Hapatitis C Virus (HCV)

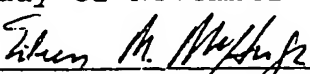
from the Italian language into the English language, and that the English text is a true and correct translation of the original, to the best of her knowledge and belief.



Elsbeth Lussi

Sworn to before me this

9th day of November 2000



EILEEN M. MEHUGH

NOTARY PUBLIC OF NEW JERSEY
My Commission Expires February 11, 2001

M E M O

DATE November 20, 2000

TO SHELDON O. HEBER LOCATION PATENT DEPT.

FROM CATHERINE KENNY LOCATION INFORMATION SERVICES
R86-210 X5830

Translation

Patent Application RM95A000343

Method for Reproducing In Vitro The RNA-
Dependent RNA Polymerase and Terminal
Nucleotidyl Transferase Activities Encoded
by Hepatitis C Virus (HCV)

R. DeFrancesco, S.E. Behrens and
L. Tomei

FOREIGN LANGUAGE: Italian

CODE: TR-2000/085

TRANSLATED BY

TRANSLATION COMPANY OF NEW YORK, INC.

FORM A
[canceled tax stamp]

TO THE MINISTER OF INDUSTRY, COMMERCE AND ARTISANRY
CENTRAL PATENTS OFFICE - ROME
PATENT APPLICATION FOR INDUSTRIAL INVENTIONS,
FILING OF RESERVATIONS AND ADVANCE PUBLIC ACCESS

A. APPLICANT (I)

Name: ISTITUTO DI RICERCHE DI BIOLOGIA MOLECOLARE P.
ANGELETTI SP
Residence: POMEZIA (ROME) ITALY I Code:
01850891001

B. REPRESENTATIVE OF APPLICANT TO THE CPO:

Last Name, First Name: de Benedetti Fabrizio et al
Name of Offices: SOCIETÀ ITALIANA BREVETTI S.p.A.
Street: Piazza di Pietra No. 0039 City: ROME
Postal Code: 00186 (Province): RM

D. TITLE

"METHOD FOR REPRODUCING IN VITRO THE RNA-DEPENDENT RNA POLYMERASE
AND TERMINAL NUCLEOTIDYL TRANSFERASE ACTIVITIES ENCODED BY
HEPATITIS C VIRUS (HCV)"

E. DESIGNATED INVENTORS

Last Name, First Name
1) DE FRANCESCO, Raffaele (Dr.)
2) BEHRENS, Sven Erik (Dr.)
3) TOMEI, Licia (Dr.)

H. SPECIAL NOTES

None

APPENDED DOCUMENTATION

	No. of Copies		
Doc. 1)	[illeg.]	No. of Pages: 54	Abstract with main drawing, description and claims (mandatory 1 copies)
Doc. 2)	1	No. of Pages: 03	Drawing (mandatory if cited in description, copies)
Doc. 3)	1		Letter of assignment
Doc. 4)	0		[illegible]

Doc. 5) 0 Designation of inventors
Doc. 6) 0 Authorization or
[illegible] of transfer
Doc. 7) 1 Complete name of
applicant

8) Certificate of payment, Total Liras Nine Hundred Fifteen Thousand, (600,000 + 315,000) mandatory

COMPILED ON 5/25/1995 SIGNATURE OF APPLICANT (1)
Gilberto Tonon
CONTINUES YES/NO YES (Registration on Roll No. 83)
[illegible signature]

EXEMPLIFIED COPY REQUESTED OF THIS CERTIFICATE Yes/No YES

PROVINCIAL OFFICE OF INDUSTRY, COMMERCE AND ARTISANRY of Rome,
RM95A000343 ROME Series 58

In the year Nineteen Hundred Ninety-Five, on the twenty-fifth day of May, the above-cited applicant(s) or the undersigned presented an application consisting of 01 pages appended hereto for issuance of the above-cited patent.

FILER ISSUING OFFICIAL
[illegible signature] [illegible signature]
[seal] [illegible stamp]
Ministry of Industry, Trade and Artisanry
Provincial Office of Industry, Trade and Artisanry

ADDITIONAL SHEET No. 01 of a Total of 01
APPLICATION RM95A000343

ADDITION, FORM A

A. APPLICANT

01 Name S.p.A.

SIGNATURE OF APPLICANT (1) *Gilberto Tonon*
(Registration on Roll No. 83)
[illegible signature]

FOR CENTRAL PATENT OFFICE USE ONLY

[seal]

Ministry of Industry, Trade and Artisanry
Provincial Office of Industry, Trade and Artisanry

SIB 90470

PROSPECTUS A

ABSTRACT, INVENTION WITH MAIN DESIGN, DESCRIPTION AND CLAIMS

APPLICATION NO.

REG. A

FILING DATE

PATENT NO.

ISSUANCE DATE

D. TITLE

"METHOD FOR REPRODUCING *IN VITRO* THE RNA-DEPENDENT RNA POLYMERASE AND TERMINAL NUCLEOTIDYL TRANSFERASE ACTIVITIES ENCODED BY HEPATITIS C VIRUS (HCV)"

L. ABSTRACT

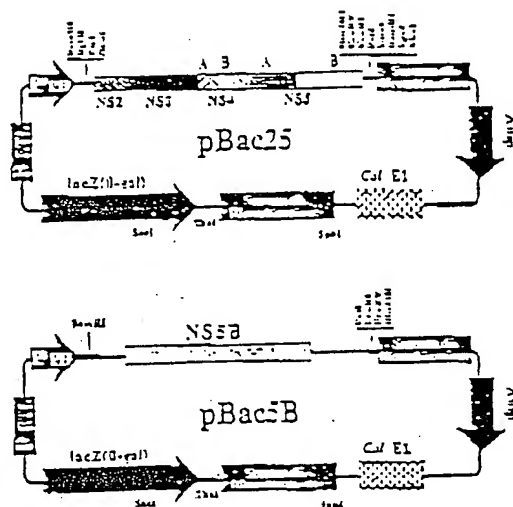
This is a method for reproducing *in vitro* the RNA-dependent RNA polymerase activity associated with hepatitis C virus. The method is characterized in that sequences contained in NS5B are used in the reaction mixture. The terminal nucleotidyl transferase activity, a further property of the NS5B protein, can also be reproduced using this method. The method takes advantage of the fact that the NS5B protein, either purified to apparent homogeneity or present in extracts of overproducing organisms, can catalyze the addition of ribonucleotides to the 3' termini of exogenous or endogenous RNA molecules.

The invention also relates to a composition of matter that comprises sequences contained in NS5B, and to the use of these compositions for the set up of an enzymatic test capable of selecting, for therapeutic purposes, compounds that inhibit the enzymatic activity associated with NS5B.

Figure 1 shows plasmids used in the method to produce hepatitis C virus RNA-dependent RNA polymerase and terminal

nucleotidyl transferase in cultivated eukaryotic and prokaryotic cells.

M. FIGURE



P_{ETL} = Promoter of the gene coding for the PCNA protein

P_{PH} = Promoter of the polyhedrin gene

Amp = Gene coding for the β-lactamase enzyme (ampicillin resistance)

LacZ (β-gal) = Gene coding for the β-galactosidase enzyme

Col E1 = pBR322 replication origin

DESCRIPTION of the Industrial Invention entitled:
"METHOD FOR REPRODUCING *IN VITRO* THE RNA-DEPENDENT RNA POLYMERASE
AND TERMINAL NUCLEOTIDYL TRANSFERASE ACTIVITIES ENCODED BY
HEPATITIS C VIRUS (HCV)"
of the Italian Institute: ISTITUTO DI RICERCHE DI BIOLOGIA
MOLECOLARE P. ANGELETTI S.p.A.
With headquarters in POMEZIA, ROME (ITALY)

DESCRIPTION

The present invention relates to the molecular biology and virology of the hepatitis C virus (HCV). More specifically, this invention has as its object the RNA-dependent RNA polymerase (RdRp) and the nucleotidyl terminal transferase (TNTase) activities produced by HCV, methods of expression of the HCV RdRp and TNTase methods to test RdRp and TNTase activities in vitro encoded by HCV in order to identify, for therapeutic purposes, compounds that inhibit these enzymatic activities and therefore might interfere with the replication of the HCV virus.

As is known, the hepatitis C virus (HCV) is the main etiological agent of hepatitis C (HCV) and the main etiological agent of non-A, non-B hepatitis (NANB). It is estimated that HCV causes at least 90% of post-transfusional NANB viral hepatitis and 50% of sporadic NANB hepatitis. Although great progress has been made in the selection of blood donors and in the immunological characterization of blood used for transfusions, there is still a high number of HCV infections among those receiving blood transfusions (one million or more infections every year throughout the world). Approximately 50% of HCV-infected individuals develop cirrhosis of the liver within a

period that can range from 5 to 40 years. Furthermore, recent clinical studies suggest that there is a correlation between chronic HCV infection and the development of hepatocellular carcinoma.

HCV is an envelope virus containing an RNA positive genome of approximately 9.4 kb. This virus is a member of the Flaviviridae family, the other members of which are the flaviviruses and the pestiviruses. The RNA genome of HCV has recently been mapped. Comparison of sequences from the HCV genomes isolated in various parts of the world has shown that these sequences can be extremely heterogeneous. The majority of the HCV genome is occupied by an open reading frame (ORF) that can vary between 9030 and 9099 nucleotides. This ORF codes for a single viral polyprotein, the length of which can vary from 3010 to 3033 amino acids. During the viral infection cycle, the polyprotein is proteolytically processed into the individual gene products necessary for replication of the virus. The genes coding for HCV structural proteins are located at the 5' end of the ORF, whereas the region coding for the non-structural proteins occupies the rest of the ORF.

The structural proteins consist of C (core, 21 kDa), E1 (envelope, gp37), and E2 (NS1, gp61). C is a non-glycosylated protein of 21 kDa that probably forms the viral nucleocapsid. The protein E1 is a glycoprotein of approximately 37 kDa, believed to be a structural protein in the outer envelope of the virus. E2, another glycoprotein with a 61 kDa membrane, probably constitutes a second structural protein of the outer envelope of the virus.

The non-structural region starts with NS2 (p24), a hydrophobic protein of 24 kDa whose function is unknown. NS3, a protein of 68 kDa that follows NS2 in the polyprotein, has two

functional domains: a serine protease domain in the first 200 amino-terminal amino acids, and an RNA-dependent ATPase domain at the carboxy terminus. The gene region corresponding to NS4 codes for NS4A (p6) and NS4B (p26), two hydrophobic proteins of 6 and 26 kDa, respectively, whose functions have not yet been clarified. The gene corresponding to NS5 also codes for two proteins, NS5A (p56) and NS5B (p65), of 56 and 65 kDa, respectively.

Various molecular biology studies indicate that the signal peptidase, a protease associated with the endoplasmic reticulum of the host cell, is responsible for proteolytic processing in the non-structural region, that is to say, at sites C/E1, E1/E2 and E2/NS2. A second protease activity of HCV appears to be responsible for the cleavage between NS2 and NS3. This protease activity is contained in a region comprising both the part of NS2 and the part of NS3 containing the serine protease domain, but does not use the same catalytic mechanism. The serine protease contained in NS3 is responsible for cleavage at the junctions between NS3 and NS4A, between NS4A and NS4B, between NS4B and NS5A, and between NS5A and NS5B.

Similarly to other (+)-strand RNA viruses, the replication of HCV is thought to proceed via the initial synthesis of a (-)-RNA strand, which serves, in turn, as template for the production of progeny (+)-strand RNA molecules. An RNA-dependent RNA polymerase (RdRp) has been postulated to be involved in both these steps. An amino acid sequence present in all the RNA-dependent RNA polymerases can be recognized within the NS5 region. This suggests that the NS5 region contains components of the viral replication machinery. Virally-encoded polymerases have traditionally been considered important targets for inhibition by antiviral compounds. In the specific case of HCV, the search for

such substances has, however, been hindered by the lack of both a suitable model system of viral infections (e.g., infection of cells in culture or a facile animal model) and a functional RdRp enzymatic assay.

It has now been unexpectedly found that this important limitation can be overcome by adopting the method according to the present invention, which also gives additional advantages that will be evident from the following.

The present invention has as its object a method for reproducing *in vitro* the RNA-dependent RNA polymerase activity of HCV that makes use of sequences contained in the HCV NS5B protein. The terminal nucleotidyl transferase activity, a further property of the NS5B protein, can also be reproduced using this method. The method takes advantage of the fact that NS5B can be expressed in either eucaryotic or prokaryotic heterologous systems: the recombinant proteins containing sequences of NS5B, either purified to apparent homogeneity or present in extracts of overproducing organisms, can catalyze the addition of ribonucleotides to the 3' termini of exogenous RNA molecules, either in a template-dependent (RdRp) or template-independent (TNTase) fashion.

The invention also extends to a new composition of matter, characterized in that it comprises proteins whose sequences are described in SEQ ID NO: 1 or sequences contained therein or derived therefrom. It is understood that this sequence may vary in different HCV isolates, since all RNA viruses show a high degree of variability. This new composition of matter has the RdRp activity necessary to HCV in order to replicate its genome.

The present invention also has as its object the use of this composition of matter in order to prepare an enzymatic assay capable of identifying, for therapeutic purposes, compounds that inhibit the enzymatic activities associated with NS5B, including inhibitors of the RdRp and that of the TNTase.

Up to this point, a general description has been given of the present invention. With the aid of the following examples, a more detailed description of specific embodiments thereof will now be given, in order to give a clearer understanding of its objects, characteristics, advantages and methods of operation.

Figure 1 shows the plasmid constructs used for HCV cDNA transfer into a baculovirus expression vector.

Figure 2 shows the plasmid used for the *in vitro* synthesis of the D-RNA substrate of the RNA-dependent RNA polymerase [pT7-7 (DCoH)], and for the expression of the RNA-dependent RNA polymerase in *E. coli* cells [pT7-7 (NS5B)], respectively.

Figure 3 shows a schematic drawing of (+) and (-)-strands of D-RNA. The transcript contains the coding region of the DCoH mRNA. The DNA-oligonucleotides a, b and c were designed to anneal with the newly-synthesized antisense RNA and the DNA/RNA hybrid was subjected to cleavage with RNase H. The lower part of the diagram depicts the expected RNA fragment sizes generated by RNase digestion of the RNA (-) hybrid with oligonucleotides a, b and c, respectively.

DEPOSITS

DH1 *E. coli* bacteria, transformed using the plasmids pBac 5B, pBac 25, pT7.7 DCoH and pT7.7NS5B - containing SEQ ID NO: 1; SEQ

ID NO: 2; the cDNA for transcription of SEQ ID NO: 12; and SEQ ID NO: 1, respectively, filed on May 9, 1995, with The National Collections of Industrial and Marine Bacteria Ltd. (NCIMB), Aberdeen, Scotland, UK, under access numbers NCIMB 40727, 40728, 40729 and 40730, respectively.

EXAMPLE 1

Method of Expression of HCV RdRp/TNTase in *Spodoptera frugiperda* Clone 9 Cultured Cells

Systems for expression of foreign genes in insect cultured cells, such as *Spodoptera frugiperda* clone 9. (Sf9) cells infected with baculovirus vectors are known in the art (V.A. Luckow, Baculovirus systems for the expression of human gene products, (1993) Current Opinion in Biotechnology 4, pp. 564 - 572). Heterologous genes are usually placed under the control of the strong polyhedrin promoter of the *Autographa californica* nuclear polyhedrosis virus of the *Bombyx mori* nuclear polyhedrosis virus. Methods for the introduction of heterologous DNA in the desired site in the baculoviral vectors by homologous recombination are also known in the art (D.R. O'Reilly, L.K. Miller, V.A. Luckow, (1992), Baculovirus Expression Vectors - A Laboratory Manual, W.H. Freeman and Company, New York).

Plasmid vectors pBac5B and pBac25 are derivatives of pBlueBacIII (Invitrogen) and were constructed for transfer of genes coding for NS5B and other non-structural HCV proteins in baculovirus expression vectors. The plasmids are schematically illustrated in Figure 1 and their construction is described in detail in Example 8. Selected fragments of the cDNA corresponding to the genome of the HCV-BK isolate (HCV-BK; Takamizawa, A., Mori, C., Fuke, I., Manabe, S., Murakami, S., Fujita, J., Onishi,

E., Andoh, T., Yoshida, I., and Okayama, H., (1991), Structure and Organization of the Hepatitis C Virus Genome Isolated from Human Carriers, *J. Virol.*, 65, 1105 - 1113) were cloned under the strong polyhedrin promoter of the nuclear polyhedrosis virus and flanked by sequences that allowed homologous recombination in a baculovirus vector.

In order to construct pBac5D, a PCR product containing the cDNA region encoding amino acids 2420 to 3010 of the HCV polyprotein and corresponding to the NS5B protein (SEQ ID NO: 1) was cloned between the *Bam*HI and *Hind*III sites of pBlue BacIII. The PCR sense oligonucleotide contained a translation initiation signal, whereas the original HCV termination codon serves for translation termination.

PBac25 is a derivative of pBlueBacIII (Invitrogen) where the cDNA region coding for amino acids 810 to 3010 of the HCV-BK polyprotein (SEQ ID NO: 2) was cloned between the *Nco*I and the *Hind*III restriction sites.

Spodoptera frugiperda clone 9 (Sf9) cells and baculovirus recombination kits were purchased from Invitrogen. Cells were grown on dishes or in suspension at 27°C in complete Grace's insect medium (Gibco) containing 10% fetal bovine serum (Gibco). Transfection, recombination, and selection of baculovirus constructs were performed as recommended by the manufacturer. Two recombinant baculovirus clones, Bac25 and Bac5B, were isolated that contained the desired HCV cDNA.

For protein expression, Sf9 cells were infected either with the recombinant baculovirus Bac25 or Bac5B at a density of 2 x 10⁶ cells per ml in a ratio of about 5 virus particles per cell. 48 - 72 hours after infection, the Sf9 cells were harvested by

centrifuging, washed once with phosphate buffered saline (PBS) and carefully resuspended (7.5×10^7 cells per ml) in buffer A (10 mM Tris/Cl pH 8, 1.5 mM MgCl₂, 10 mM NaCl) containing 1 mM dithiothreitol (DTT), 1 mM phenylmethylsulfonyl fluoride (PMSF, Sigma), and 4 mg/ml leupeptin. All the following steps were performed between 0° and 4°C: after swelling for 30 minutes, the cells were disrupted by 20 strokes in a Dounce homogenizer using a tight-fitting pestle. Glycerol, as well as the detergents Nonidet P-40 (NP40) and 3-[(3-Cholamidopropyl)-dimethyl-ammonio]-1-propanesulfonate (CHAPS), were added to final concentrations of 10% (v/v), 1% (v/v) and 0.5% (w/v), respectively, and the cellular extract was incubated for a further hour on ice with occasional agitation. The nuclei were pelleted by centrifugation for 10 minutes at 1000 x g, and the supernatant was collected. The pellet was resuspended in buffer A containing the above concentrations of glycerol and detergents (0.5 ml per 7.5×10^7 nuclei) by 20 strokes in the Dounce homogenizer and then incubated for one hour on ice. After repelleting the nuclei, both supernatants were combined, centrifuged for 10 minutes at 8000 x g and the pellet was discarded. The resulting crude cytoplasmic extract was used either directly to determine the RdRp activity or further purified on a sucrose gradient (see Example 5).

Infection of Sf9 cells with either the recombinant baculovirus Bac25 or Bac5B leads to the expression of the expected HCV proteins. Indeed, following infection of Sf9 cells with Bac25, correctly-processed HCV NS2 (24 kDa), NS3 (68 kDa), NS4B (26 kDa), NS4A (6 kDa), NS5A (56 kDa), and NS5B (65 kDa) proteins can be detected in the cell lysates by SDS polyacrylamide gel electrophoresis (SDS-PAGE) and immunostaining. Following infection of Sf9 cells with Bac5B, only one HCV-encoded protein, corresponding in size to authentic NS5B (65 kDa), is

detected by SDS-PAGE followed by immuno- or Coomassie Blue staining.

EXAMPLE 2

Method of Assay of Recombinant HCV RdRp on a Synthetic RNA Template/Substrate

The RdRp assay is based on the detection of labeled nucleotides incorporated into novel RNA products. The *in vitro* assay to determine RdRp activity was performed in a total volume of 40 μ l containing 1 - 5 μ l of either Sf9 crude cytoplasmic extract or purified protein fraction. Unfractionated or purified cytoplasmic Sf9 cell extracts infected with Bac25 or Bac5D may be used as the source of HCV RdRp. A Sf9 cell extract obtained from cells infected with a recombinant baculovirus construct expressing a protein that is not related to HCV may be used as a negative control. The following supplements are added to the reaction mixture (final concentrations): 20 mM Tris/Cl pH 7.5, 5 mM MgCl₂, 1 mM DTT, 25 mM KCl, 1 mM EDTA, 5 - 10 μ Ci [³²P] NTP of one species (unless otherwise specified, GTP, 3000 Ci/mmol, Amersham, was used), 0.5 mM each NTP (i.e., CTP, UTP, ATP unless otherwise specified), 20 U RNasin (Promega), 0.5 μ g RNA-substrate (ca. 4 pmol; final concentration 100 nM), 2 μ g actinomycin D (Sigma). The reaction was incubated for two hours at room temperature, stopped by the addition of an equal volume of 2 x Proteinase K (PK, Boehringer Mannheim) buffer (300 mM NaCl, 100 mM Tris/Cl pH 7.5, 1% w/v SDS) and followed by half an hour of treatment with 50 μ g of PK at 37°C. RNA products were PCA extracted, precipitated with ethanol, and analyzed by electrophoresis on 5% polyacrylamide gels containing 7M urea.

The RNA substrate we normally used for the assay (D-RNA) had the sequence reported in SEQ ID NO: 12, and was typically obtained by *in vitro* transcription of the plasmid pT7-7 (DCoH) with T7 polymerase, as described below.

Plasmid pT7-7 (DCoH) (Figure 2) was linearized with the unique *Bgl*III restriction site contained at the end of the DCoH coding sequence and transcribed *in vitro* with T7 polymerase (Stratagene) using the procedure described by the manufacturer. Transcription was stopped by the addition of 5 U/10 μ l of DnaseI (Promega). The mixture was incubated for a further 15 minutes and extracted with phenol/chloroform/isoamyl alcohol (PCA). Unincorporated nucleotides were removed by gel filtration through a 1 ml Sphadex G50 spun column. After extraction with PCA and ethanol precipitation, the RNA was dried, redissolved in water, and its concentration determined by optical density at 260 nm.

As will be clear from the experiments described below, any RNA molecule other than D-RNA may be used for the RdRp assay of the invention.

The above-described HCV RdRp assay gave rise to a characteristic pattern of radioactively-labeled reaction products: one labeled product that co-migrated with the substrate RNA was observed in all reactions, including the negative control. This RNA species could also be visualized by silver staining and was thus thought to correspond to the input substrate RNA, labeled most likely by terminal nucleotidyl transferase activities present in cytoplasmic extracts of baculovirus-infected Sf9 cells. In the reactions carried out with the cytoplasmic extracts of Sf9 cells infected with either Bac25 or Bac5B, but not of cells infected with a recombinant baculovirus construct expressing a protein that is not related to

HCV, an additional band was observed, migrating faster than the substrate RNA. This latter reaction product was found to be labeled to a high specific activity, since it could be detected solely by autoradiography and not by silver staining. This novel product was found to be derived from the externally-added RNA template, as it was absent from control reactions where no RNA was added. Interestingly, the formation of a labeled species migrating faster than the substrate RNA was consistently observed with a variety of template RNA molecules, whether containing the HCV 3' untranslated region or not. The messenger RNA of the liver-specific transcription cofactor DCoH (D-RNA) turned out to be an efficiently accepted substrate in our RdRp assay.

In order to define the nature of the species generated in the reaction by the Bac25- or Bac5B-infected cell extracts, we carried out the following series of experiments: (i) The product mixture was treated with RNase A or Nuclease P1. As this resulted in the complete disappearance of the radioactive bands, we concluded that both the labeled products were RNA molecules. (ii) Omission from the reaction mixtures of any of the four nucleotide triphosphates resulted in labeling of only the input RNA, suggesting that the faster-migrating species is a product of a polymerization reaction. (iii) Omission of Mg^{2+} ions from the assay caused a complete block of the reaction: neither synthesis of the novel RNA nor labeling of the RNA substrate was observed. (iv) When the assay was carried out with a radioactively labeled input RNA and unlabeled nucleotides, the labeled product was indistinguishable from that obtained under the standard conditions. We concluded from this result that the novel RNA product is generated from the original input RNA molecule.

Taken together, our data demonstrate that the extracts of Bac25- or Bac5B-infected Sf9 cells contain a novel magnesium-

dependent enzymatic activity that catalyzes *de novo* RNA synthesis. This activity was shown to be dependent upon the presence of added RNA, but independent of an added primer or of the origin of the input RNA molecule. Moreover, as the products generated by extracts of Sf9 cells infected with either Bac25 or Bac5B appeared to be identical, the experiments just described indicate that the observed RdRp activity is encoded by the HCV NS5B protein.

EXAMPLE 3

Methods for the Characterization of the HCV RdRp RNA Product

The following methods were employed in order to elucidate the structural RNA product. Under our standard electrophoresis conditions (5% polyacrylamide, 7M urea), the size of the novel RNA product appeared to be approximately 200 nucleotides. This could be due to either internal initiation of RNA transcription, or to premature termination. These possibilities, however, appeared to be very unlikely, since products derived from RdRp assays using different RNA substrates were all found to migrate significantly faster than their respective templates. Increasing the temperature during electrophoresis and the concentration of acrylamide in the analytic gel leads to a significantly different migration behavior of the RdRp product. Thus, using, for instance, a gel system containing 10% acrylamide, 7M urea, where separation was carried out at higher temperatures, the RdRp product migrated slower than the input substrate RNA, at a position corresponding to at least double the length of the input RNA. A similar effect was observed when RNA-denaturing agents such as methylhydroxymercury (CH_3HgOH , 10 mM) were added to the RdRp products prior to electrophoresis on a low-percentage/lower

temperature gel. These observations suggest that the RdRp product possesses an extensive secondary structure.

We investigated the susceptibility of the product molecule to a variety of ribonucleases of different specificity. The product was completely degraded upon treatment with RNase A. On the other hand, it was found to be surprisingly resistant to single-strand specific nuclease RNase T1. The input RNA was completely degraded after 10 minutes incubation with 60 U RNase T1 at 22°C, and silver staining of the same gel confirmed that not only the template, but also all other RNA usually detectable in the cytoplasmic extracts of Sf9 cells, was completely hydrolyzed during incubation with RNase T1. In contrast, the RdRp product remained unaltered and was affected only following prolonged incubation with RNase T1. Thus, after 2 hours of treatment with RNase T1 the labeled product molecule could no longer be detected at its original position in the gel. Instead, a new band appeared that had an electrophoretic mobility similar to the input template RNA. A similar effect was observed when carrying out the RNase T1 digestion for 1 hour, but at different temperatures: at 22°C, the RdRp product remained largely unaffected whereas at 37°C it was converted to the new product that co-migrated with the original substrate.

The explanation for these observations is that the input RNA serves as a template for the HCV RdRp, where the 3'-OH is used to prime the synthesis of the complementary strand by a "turn" or "copy-back" mechanism to give rise to a duplex RNA "hairpin" molecule, consisting of the sense (template) strand to which an antisense strand is covalently attached. Such a structure would explain the unusual electrophoretic mobility of the RdRp product on the polyacrylamide gels as well as its high resistance to single-strand specific nucleases. The hairpin RNA molecule should

not be base-paired and therefore ought to be accessible to the nucleases. Treatment with RNase T1 thus leads to the hydrolysis of the covalent link between the sense and antisense strands to yield a double-stranded RNA molecule. During denaturing gel electrophoresis, the two strands become separated and only the newly-synthesized antisense strand, which should be similar in length to the original RNA template, would remain detectable. This mechanism would appear rather likely, especially in view of the fact that this kind of product is generated by several other RNA polymerases *in vitro*.

The following experiment was designed to demonstrate that the RNA product labeled during the polymerase reaction and apparently released by RNase T1 treatment exhibits antisense orientation with respect to the input template.

For this purpose, we synthesized oligodeoxyribonucleotides corresponding to three separate sequences of the input template RNA molecule (Figure 2), oligonucleotide a, corresponding to nucleotides 170 - 195 of D-RNA (SEQ ID NO: 3); oligonucleotide b, complementary to nucleotides 286 - 309 (SEQ ID NO: 4); oligonucleotide c, complementary to nucleotides 331 - 354 (SEQ ID NO: 5). These were used to generate DNA/RNA hybrids with the product of the polymerase reaction, such that they could be subjected to RNase H digests. Initially, the complete RdRp product was used in the hybridizations. However, as this structure is too thermostable, no specific hybrids were formed. The hairpin RNA was therefore pre-treated with RNase T1, denatured by boiling for 5 minutes and then allowed to cool down to room temperature in the presence of the respective oligonucleotide. As expected, exposure of the hybrids to RNase H yielded specific cleavage products. Oligonucleotide a-directed cleavage led to products of about 170 and 220 nucleotides in

length, oligonucleotide b yielded products of about 290 and 110 nucleotides and oligonucleotide c gave rise to fragments of about 330 and 65 nucleotides. As these fragments are of the expected sizes (see Figure 3), the results indicate that the HCV NS5B-mediated RNA synthesis proceeds by a copy-back mechanism that generates a hairpin-like RNA duplex.

EXAMPLE 4

Method of Assay of Recombinant HCV TNTase on a Synthetic RNA Substrate

The TNTase assay is based on the detection of template-independent incorporation of labeled nucleotides to the 3' hydroxyl group of RNA substrates. The DNA substrate for the assay (D-RNA) was typically obtained by *in vitro* transcription of the linearized plasmid pT7-7DCoH with T7 polymerase as described in Example 2. However, any RNA molecule other than D-RNA may be used for the TNTase assay of the invention.

The *in vitro* assay to determine TNTase activity was performed in a total volume of 40 μ l containing 1 - 5 μ l of either Sf9 crude cytoplasmic extract of purified protein fraction. Unfractionated or purified cytoplasmic extracts of Sf9 cells infected with Bac25 or Bac5B may be used as the source of HCV TNTase. An Sf9 cell extract obtained from cells infected with a recombinant baculovirus construct expressing a protein that is not related to HCV may be used as a negative control. The following supplements are added to the reaction mixture (final concentrations): 20 mM Tris/Cl pH 7.5, 5 mM MgCl₂, 1 mM DTT, 25 mM KCl, 1 mM EDTA, 5 - 10 μ Ci [³²P] NTP of one species (unless otherwise specified, UTP, 3000 Ci/mmol, Amersham, was used), 20 U RNasin (Promega), 0.5 μ g RNA-substrate (ca. 4 pmol); final

concentration 100 mM), 2 µg actinomycin D (Sigma). The reaction was incubated for two hours at room temperature, stopped by the addition of an equal volume of 2 x Proteinase K (PK, Boehringer Mannheim) buffer (300 mM NaCl, 100 mM Tris/Cl pH 7.5, 1% w/v SDS) and followed by half an hour of treatment with 50 µg of PK at 37°C. RNA products were PCA extracted, precipitated with ethanol and analyzed by electrophoresis on 5% polyacrylamide gels containing 7M urea.

EXAMPLE 5

Method for the Purification of the HCV RdRp/TNTase by Sucrose Gradient Sedimentation

A linear 0.3 – 1.5 M sucrose gradient was prepared in buffer A containing detergents (see Example 1). Up to 2 ml of extract of Sf9 cells infected with Bac5B or Bac25 (corresponding to about 8×10^7 cells) were loaded onto a 12 ml gradient. Centrifugation was carried out for 20 hours at 39000 x g using a Beckmann SW40 rotor. 0.5 ml fractions were collected and assayed for activity. The NS5B protein, identified by immunostaining, was found to migrate in the density gradients with an unexpectedly high sedimentation coefficient. The viral protein and ribosomes were found to co-sediment in the same gradient fractions. This behavior enabled us to separate the viral protein from the main bulk of cytoplasmic proteins, which remained on the top of the gradient. The RdRp activity assay revealed that the RdRp activity co-sedimented with the NS5B protein. A terminal nucleotidyl transferase activity was also present in these fractions.

EXAMPLE 6

Method for the Purification of the HCV TNTase/RdRp from Sf9 Cells

Whole cell extracts are made from 1 g of Sf9 cells infected with Bac5B recombinant baculovirus. The frozen cells are thawed on ice in 10 ml of buffer containing 20 mM Tris/HCl pH 7.5, 1 mM EDTA, 10 mM DTT, 50% glycerol (N buffer) supplemented with 1 mM PMSF. Triton X-100 and NaCl are then added to a final concentration of 2% and 500 mM, respectively, in order to promote cell breakage. After the addition of MgCl₂ (10 mM) and DNase I (15 µg/ml), the mixture is stirred at room temperature for 30 minutes. The extract is then cleared by ultracentrifugation in a Beckman centrifuge, using a 90 Ti rotor at 40000 rpm for 30 minutes at 4°C. The extract is diluted with a buffer containing 20 mM Tris/HCl pH 7.5, 1 mM EDTA, 10 mM DTT, 20% glycerol, 0.5% Triton X-100 (LG buffer) in order to adjust the NaCl concentration to 300 mM, and incubated with 5 ml of DEAE-Sepharose Fast Flow, equilibrated in LG buffer containing 300 mM NaCl. The matrix is then poured into a column and washed with two volumes of the same buffer. The "flow-through" and the first wash of the DEAE-Sepharose Fast Flow column is diluted 1:3 with LG buffer and applied onto a heparin-Sepharose CL6B column (10 ml) equilibrated with LG buffer containing 100 mM NaCl. The Heparin-Sepharose CL6B is washed thoroughly and the bound proteins are eluted with a linear 100 ml gradient, from 100 mM to 1M NaCl in LG buffer. The fractions containing NS5B, as judged by silver- and immunostaining of SDS-PAGE, are pooled and diluted with LG buffer in order to adjust the NaCl concentration to 50 mM. The diluted fractions are subsequently applied to a Mono Q-FPLC column (1 ml) equilibrated with LG buffer containing 50 mM NaCl. Proteins are eluted with a linear gradient (20 ml) from 50 mM to 1 M NaCl in LG buffer. The fractions containing NS5B, as judged by silver- and immunostaining of SDS-PAGE, are pooled and diluted with LG buffer to adjust the NaCl concentration to 50 mM. The fractions containing NS5B, as judged by silver- and

immunostaining by SDS-PAGE, are pooled and dialyzed against LG buffer containing 100 mM NaCl. After extensive dialysis, the pooled fractions were loaded onto poly(U)-Sepharose CL6B (10 ml) equilibrated with LG buffer containing 100 mM NaCl. The poly(U)-Sepharose CL6B was washed thoroughly and the bound proteins were eluted with a linear 100 ml gradient, from 100 mM to 1M NaCl in LG buffer. The fractions containing NS5B, as judged by silver- and immunostaining of SDS-PAGE, are pooled, dialyzed against LG buffer containing 100 mM NaCl and stored in liquid nitrogen prior to activity assay.

Fractions containing the purified protein NS5B were tested for the presence of both activities. The RdRp and TNTase activities were found in the same fractions. These results indicate that both activities, RNA-dependent RNA polymerase and terminal ribonucleotide transferase, are the functions of the HCV NS5B protein.

We tested the purified NS5B for terminal nucleotidyl transferase activity with each of the four ribonucleotide triphosphates at non-saturating substrate concentrations. The results clearly showed that UTP is the preferred TNTase substrate, followed by ATP, CTP and GTP, regardless of the origin of the input RNA.

EXAMPLE 7

Method of Expression of HCV RdRp/TNTase in E. coli

The plasmid pT7-7(NS5B), described in Figure 2 and Example 8, was constructed in order to allow expression in E. coli of the HCV protein fragment having the sequence reported in SEQ ID NO 1. Such protein fragment contains the RdRp and the TNTase of NS5B,

as discussed above. The fragment of HCV cDNA coding for the NS5B protein was thus cloned downstream of the bacteriophage T7 Ø10 promoter and in frame with the first ATG codon of the phage T7 gene 10 protein, using methods that are known in molecular biology practice and described in detail in Example 8. The pT7-7(NS5B) plasmid also contains the gene for the β -lactamase enzyme that can be used as a marker of selection of *E. coli* cells transformed with plasmid pT7-7(NS5B).

The plasmid pT7-7(NS5B) was then transformed in the *E. coli* strain BL21(DE53), which is normally employed for high-level expression of genes cloned into expression vectors containing T7 promoter. In this strain of *E. coli*, the T7 gene polymerase is carried on the bacteriophage λ DE53, which is integrated into the chromosome of BL21 cells (Studier and Moffatt, Use of bacteriophage T7 RNA polymerase to direct selective high-level expression of cloned genes, (1986), *J. Mol. Biol.* 189, pp. 113 - 130). Expression from the gene of interest is induced by addition of isopropylthiogalactoside (IPTG) to the growth medium according to a procedure that has been previously described (Studier and Moffatt, 1986). The recombinant NS5B protein fragment containing the RdRp is thus produced in the inclusion bodies of the host cells. Recombinant NS5B protein can be purified from the particulate fraction of *E. coli* BL21(DE53) extracts and refolded according to procedures that are known in the art (D.R. Thatcher and A. Hitchcock, Protein folding in Biotechnology, (1994), in "Mechanism of protein folding", R.H. Pain, EDITOR, IRL PRESS, pp. 229 - 255).

EXAMPLE 8

Detailed Construction of the Recombinant Vectors Expressing NS5B

Selected fragments of the cDNA corresponding to the genome of the HCV-BK isolate (HCVBK) were cloned under the strong polyhedrin promoter of the nuclear polyhedrosis virus and flanked by sequences that allowed homologous recombination in a baculovirus vector.

pBac5B contains the HCV-BK sequence comprised between nucleotide 7590 and 9366, and codes for the NS5B protein reported in SEQ ID NO: 1. In order to obtain this plasmid, a cDNA fragment was generated by PCR using synthetic oligonucleotides having the sequences 5'-AAGGATCCATGTCAATGTCCCTACACATGGAC-3' (SEQ ID NO: 6) and 5'-AATATTGAAATTCATCGGTTGGGGAGCAGGTAGATG-3' (SEQ ID NO: 7), respectively. The PCR product was then treated with the Klenow DNA polymerase, digested at the 5' end with *BamHI*, and subsequently cloned between the *BamHI* and *SmaI* sites of the Bluescript SK(+) vector. Subsequently, the cDNA fragment of interest was digested out with the restriction enzymes *BamHI* and *HindIII* and religated in the same sites of the pBlueBacIII vector (Invitrogen).

pBac25 contains the HCV-BK cDNA region comprised between nucleotides 2759 and 9416 and codes for amino acids 810 to 3010 of the HCV-BK polyprotein (SEQ ID NO: 2). This construct was obtained as follows: first, the 820 bp cDNA fragment containing the HCV-BK sequence comprised between nucleotides 2759 and 3578 was obtained from pCD(38-9.4) (Tomei, L., Failla, C., Santolini, E., De Francesco, R., and La Monica, N. (1993) NS3 is a Serine Protease Required for Processing of Hepatitis C Virus Polyprotein, *J. Virol.*, 67, 4017 - 4026) by digestion with *NcoI* and cloned in the *NcoI* site of the pBlueBacIII vector (Invitrogen) yielding a plasmid called pBacNCO. The cDNA fragment containing the HCV-BK sequence comprised between nucleotides 1959 and 9416 was obtained from pCD(38-9.4) (Tomei et al, 1993) by

digestion with *NotI* and *XbaI* and cloned in the same sites of the Bluescript SK(+) vector yielding a plasmid called pB1sNX. The cDNA fragment containing the HCV-BK sequence comprised between nucleotides 3304 and 9416 was obtained from pB1sNX by digestion with *SacII* and *HindIII* and cloned in the same sites of the pB1sNX plasmid, yielding the pBac25 plasmid.

PT7-7 (DCoH) contains the entire coding region (316 nucleotides) of the rat dimerization cofactor of hepatocyte nuclear factor-1 α (DCoH; Mendel, D.B., Khavari, P.A., Conley, P.B., Graves, M.K., Hansen, L.P., Admon, A., and Crabtree, G.R., (1991), Characterization of a Cofactor that Regulates Dimerization of a Mammalian Homeodomain Protein, *Science*, 254, 1762 - 1767; GenBank accession number: M83740). The cDNA fragment corresponding to the coding sequence for rat DCoH was amplified by PCR using the synthetic oligonucleotide Dpr1 and Dpr2 that have the sequence TGGCTGGCAAGGCACACAGGCT (SEQ ID NO.: 8) and AGGCAGGGTAGATCTATGTC (SEQ ID NO: 9), respectively. The cDNA fragment thus obtained was cloned into the *SmaI* restriction site of the *E. coli* expression vector pT7-7. The pT7-7 expression vector is a derivative of pBR322 that contains, in addition to the β -lactamase gene and the *Col E1* origin of replication, the T7 polymerase promoter $\emptyset 10$ and the translational start site for the T7 gene 10 protein (Tabor, S., and Richardsonson, C.C. (1985) A bacteriophage T7 RNA polymerase/promoter system for controlled exclusive expression of specific genes, *Proc. Natl. Acad. Sci. USA*, 82, 1074 - 1078).

pT7-7 (NS5B) contains the HCV sequence from nucleotide 7590 to nucleotide 9366, and codes for the NS5B protein reported in SEQ ID NO: 1. In order to obtain this plasmid, a cDNA fragment was generated by PCR using oligonucleotides having the sequences 5'-TCAATGTCCTACACATGGGAC-3' (SEQ ID NO: 10) and 5' -

GATCTCTAGATCATCGGTTGGGGAGGAGGTAGATGCC-3' (SEQ ID NO: 11), respectively. The PCR product was then treated with the Klenow DNA polymerase, and subsequently ligated in the *E. coli* expression vector pT7-7 after linearizing it with *EcoRI* and blunting its extremities with the Klenow DNA polymerase.

SEQUENCE LISTING

GENERAL INFORMATION

(i) APPLICANT: ISTITUTO DI RICERCHE DI BIOLOGIA MOLECOLARE P. ANGELETTI S.p.A.

(ii) TITLE OF INVENTION: METHOD FOR REPRODUCING IN VITRO THE RNA-DEPENDENT RNA POLYMERASE AND TERMINAL NUCLEOTIDYL TRANSFERASE ACTIVITIES ENCODED BY HEPATITIS C VIRUS (HCV)

(iii) NUMBER OF SEQUENCES: 12

(iv) CORRESPONDENCE ADDRESS:

(A) ADDRESSEE: Società Italiana Brevetti

(B) STREET: Piazza di Pietra, 39

(C) CITY: Rome

(D) COUNTRY: Italy

(E) POSTAL CODE: 1-00186

(v) COMPUTER READABLE FORM:

(A) MEDIUM TYPE: Floppy Disk 3.5" 1.44 MB

(B) COMPUTER: IBM PC Compatible

(C) OPERATING SYSTEM: PC-DOS/MS-DOS Rev. 5.0

(D) SOFTWARE: Microsoft Word 6.0

(viii) ATTORNEY INFORMATION

(A) NAME: DI CERBO, Mario (Dr.)

(C) REFERENCE: RM/X90740/IN-DC

(ix) TELECOMMUNICATION INFORMATION

(A) TELEPHONE: 06/6785941

(B) TELECOPIER: 06/6794692

(C) TELEX: 612287 ROPAT

(1) INFORMATION FOR SEQ ID NO: 1:

(i) SEQUENCE CHARACTERISTICS

(A) LENGTH: 591 amino acids

(B) TYPE: amino acid

(C) NUMBER OF CHAINS: single
(D) CONFIGURATION: linear
(ii) MOLECULE TYPE: protein
(iii) HYPOTHETICAL: No
(iv) ANTISENSE: No
(v) FRAGMENT TYPE: C-terminal fragment
(vi) ORIGINAL SOURCE:
(A) ORGANISM: Hepatitis C Virus
(C) ISOLATE: BK
(vii) IMMEDIATE SOURCE: cDNA clone pCD(38-9.4) described
by Tomei et al, 1993
(ix) CHARACTERISTICS:
(A) NAME: NS5B Non-structural protein
(C) IDENTIFICATION METHOD: Experimental
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1

[SEQUENCES]

-38-

Asn Cys Gly Tyr Arg Arg Cys Arg Ala Ser Gly Val Leu Thr Thr Ser
 275 280 285
 Cys Gly Asn Thr Leu Thr Cys Tyr Leu Lys Ala Ser Ala Ala Cys Arg
 290 295 300
 Ala Ala Lys Leu Gln Asp Cys Thr Met Leu Val Asn Gly Asp Asp Leu
 305 310 315 320
 Val Val Ile Cys Glu Ser Ala Gly Thr Gln Glu Asp Ala Ala Ser Leu
 325 330 335
 Arg Val Phe Thr Glu Ala Met Thr Arg Tyr Ser Ala Pro Pro Gly Asp
 340 345 350
 Pro Pro Gln Pro Glu Tyr Asp Leu Glu Leu Ile Thr Ser Cys Ser Ser
 355 360 365
 Asn Val Ser Val Ala His Asp Ala Ser Gly Lys Arg Val Tyr Tyr Leu
 370 375 380
 Thr Arg Asp Pro Thr Thr Pro Leu Ala Arg Ala Ala Trp Glu Thr Ala
 385 390 395 400
 Arg His Thr Pro Val Asn Ser Trp Leu Gly Asn Ile Ile Met Tyr Ala
 405 410 415
 Pro Thr Leu Trp Ala Arg Met Ile Leu Met Thr His Phe Phe Ser Ile
 420 425 430
 Leu Leu Ala Gln Glu Gln Leu Glu Lys Ala Leu Asp Cys Gln Ile Tyr
 435 440 445
 Gly Ala Cys Tyr Ser Ile Glu Pro Leu Asp Leu Pro Gln Ile Ile Glu
 450 455 460
 Arg Leu His Gly Leu Ser Ala Phe Ser Leu His Ser Tyr Ser Pro Gly
 465 470 475 480
 Glu Ile Asn Arg Val Ala Ser Cys Leu Arg Lys Leu Gly Val Pro Pro
 485 490 495
 Leu Arg Val Trp Arg His Arg Ala Arg Ser Val Arg Ala Arg Leu Leu
 500 505 510
 Ser Gln Gly Gly Arg Ala Ala Thr Cys Gly Lys Tyr Leu Phe Asn Trp
 515 520 525
 Ala Val Lys Thr Lys Leu Lys Leu Thr Pro Ile Pro Ala Ala Ser Arg
 530 535 540
 Leu Asp Leu Ser Gly Trp Phe Val Ala Gly Tyr Ser Gly Gly Asp Ile
 545 550 555 560
 Tyr His Ser Leu Ser Arg Ala Arg Pro Arg Trp Phe Met Leu Cys Leu
 565 570 575
 Leu Leu Leu Ser Val Gly Val Gly Ile Tyr Leu Leu Pro Asn Arg
 580 585 590

(2) INFORMATION FOR SEQ ID NO: 2:

(i) SEQUENCE CHARACTERISTICS

- (A) LENGTH: 2201 amino acids
- (B) TYPE: amino acid
- (C) NUMBER OF CHAINS: single
- (D) CONFIGURATION: linear

(ii) MOLECULE TYPE: polypeptide

(iii) HYPOTHETICAL: No

(iv) ANTISENSE: No

(v) FRAGMENT TYPE: C-terminal fragment

(vii) IMMEDIATE SOURCE: cDNA clone pCD(38-9.4) described
by Tomei et al, 1993

(ix) CHARACTERISTICS:

- (A) NAME: NS2-NS5B Non-structural protein
precursor

(C) IDENTIFICATION METHOD: Experimental

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2

[SEQUENCES]

-39-

(2) INFORMAZIONI SULLA SEQ ID NO: 2:

(i) CARATTERISTICHE DELLA SEQUENZA:

- (A) LUNGHEZZA: 2201 amminoacidi
- (B) TIPO: amminoacididica
- (C) NUMERO DI CATENE: singola
- (D) CONFIGURAZIONE: lineare

(ii) TIPO DI MOLECOLA: polipeptide

(iii) IPOTETICA: No

(iv) ANTISENSO: No

(v) TIPO DI FRAMMENTO: frammento C-terminale

(vi) SORGENTE IMMEDIATA: cDNA clone pCD(36-9.4) descritto da Tomei ed altri, 1993

(ix) CARATTERISTICHE:

- (A) NOME: NS2-NS5B Nonstructural Protein Precursor
- (C) METODO DI IDENTIFICAZIONE: Sperimentale

(xi) DESCRIZIONE DELLA SEQUENZA: SEQ ID NO: 2:

Met Asp Arg Glu Met Ala Ala Ser Cys Gly Gly Ala Val Phe Val Gly
 1 5 10 15

Leu Val Leu Leu Thr Leu Ser Pro Tyr Tyr Lys Val Phe Leu Ala Arg
 20 25 30

Leu Ile Trp Trp Leu Gln Tyr Phe Thr Thr Arg Ala Glu Ala Asp Leu
 35 40 45

His Val Trp Ile Pro Pro Leu Asn Ala Arg Gly Arg Asp Ala Ile
 50 55 60

Ile Leu Leu Met Cys Ala Val His Pro Glu Leu Ile Phe Asp Ile Thr
 65 70 75 80

Lys Leu Leu Ile Ala Ile Leu Gly Pro Leu Met Val Leu Gln Ala Gly
 85 90 95

Ile Thr Arg Val Pro Tyr Phe Val Arg Ala Gln Gly Leu Ile His Ala
 100 105 110

Cys Met Leu Val Arg Lys Val Ala Gly Gly His Tyr Val Gln Met Ala
 115 120 125

Phe Met Lys Leu Gly Ala Leu Thr Gly Thr Tyr Ile Tyr Asn His Leu
 130 135 140

Thr Pro Leu Arg Asp Trp Pro Arg Ala Gly Leu Arg Asp Leu Ala Val
 145 150 155 160

Ala Val Glu Pro Val Val Phe Ser Asp Met Glu Thr Lys Ile Ile Thr
 165 170 175

Trp Gly Ala Asp Thr Ala Ala Cys Gly Asp Ile Ile Leu Gly Leu Pro
 180 185 190

-40-

Val Ser Ala Arg Arg Gly Lys Glu Ile Leu Leu Gly Pro Ala Asp Ser
 195 200 205
 Leu Glu Gly Arg Gly Leu Arg Leu Leu Ala Pro Ile Thr Ala Tyr Ser
 210 215 220
 Gln Gln Thr Arg Gly Leu Leu Gly Cys Ile Ile Thr Ser Leu Thr Gly
 225 230 235 240
 Arg Asp Lys Asn Gln Val Glu Gly Glu Val Gln Val Val Ser Thr Ala
 245 250 255
 Thr Gln Ser Phe Leu Ala Thr Cys Val Asn Gly Val Cys Trp Thr Val
 260 265 270
 Tyr His Gly Ala Gly Ser Lys Thr Leu Ala Ala Pro Lys Gly Pro Ile
 275 280 285
 Thr Gln Met Tyr Thr Asn Val Asp Gln Asp Leu Val Gly Trp Pro Lys
 290 295 300
 Pro Pro Gly Ala Arg Ser Leu Thr Pro Cys Thr Cys Gly Ser Ser Asp
 305 310 315 320
 Leu Tyr Leu Val Thr Arg His Ala Asp Val Ile Pro Val Arg Arg Arg
 325 330 335
 Gly Asp Ser Arg Gly Ser Leu Leu Ser Pro Arg Pro Val Ser Tyr Leu
 340 345 350
 Lys Gly Ser Ser Gly Gly Pro Leu Leu Cys Pro Phe Gly His Ala Val
 355 360 365
 Gly Ile Phe Arg Ala Ala Val Cys Thr Arg Gly Val Ala Lys Ala Val
 370 375 380
 Asp Phe Val Pro Val Glu Ser Met Glu Thr Thr Met Arg Ser Pro Val
 385 390 395 400
 Phe Thr Asp Asn Ser Ser Pro Pro Ala Val Pro Gln Ser Phe Gln Val
 405 410 415
 Ala His Leu His Ala Pro Thr Gly Ser Gly Lys Ser Thr Lys Val Pro
 420 425 430
 Ala Ala Tyr Ala Ala Gln Gly Tyr Lys Val Leu Val Leu Asn Pro Ser
 435 440 445
 Val Ala Ala Thr Leu Gly Phe Gly Ala Tyr Met Ser Lys Ala His Gly
 450 455 460
 Ile Asp Pro Asn Ile Arg Thr Gly Val Arg Thr Ile Thr Thr Gly Ala
 465 470 475 480
 Pro Val Thr Tyr Ser Thr Tyr Gly Lys Phe Leu Ala Asp Gly Gly Cys
 485 490 495
 Ser Gly Gly Ala Tyr Asp Ile Ile Ile Cys Asp Glu Cys His Ser Thr
 500 505 510

-41-

Asp Ser Thr Thr Ile Leu Gly Ile Gly Thr Val Leu Asp Gln Ala Glu
 515 520 525
 Thr Ala Gly Ala Arg Leu Val Val Leu Ala Thr Ala Thr Pro Pro Gly
 530 535 540
 Ser Val Thr Val Pro His Pro Asn Ile Glu Glu Val Ala Leu Ser Asn
 545 550 555 560
 Thr Gly Glu Ile Pro Phe Tyr Gly Lys Ala Ile Pro Ile Glu Ala Ile
 565 570 575
 Arg Gly Gly Arg His Leu Ile Phe Cys His Ser Lys Lys Lys Cys Asp
 580 585 590
 Glu Leu Ala Ala Lys Leu Ser Gly Leu Gly Ile Asn Ala Val Ala Tyr
 595 600 605
 Tyr Arg Gly Leu Asp Val Ser Val Ile Pro Thr Ile Gly Asp Val Val
 610 615 620
 Val Val Ala Thr Asp Ala Leu Met Thr Gly Tyr Thr Gly Asp Phe Asp
 625 630 635 640
 Ser Val Ile Asp Cys Asn Thr Cys Val Thr Gln Thr Val Asp Phe Ser
 645 650 655
 Leu Asp Pro Thr Phe Thr Ile Glu Thr Thr Val Pro Gln Asp Ala
 660 665 670
 Val Ser Arg Ser Gln Arg Arg Gly Arg Thr Gly Arg Gly Arg Arg Gly
 675 680 685
 Ile Tyr Arg Phe Val Thr Pro Gly Glu Arg Pro Ser Gly Met Phe Asp
 690 695 700
 Ser Ser Val Leu Cys Glu Cys Tyr Asp Ala Gly Cys Ala Trp Tyr Glu
 705 710 715 720
 Leu Thr Pro Ala Glu Thr Ser Val Arg Leu Arg Ala Tyr Leu Asn Thr
 725 730 735
 Pro Gly Leu Pro Val Cys Gln Asp His Leu Glu Phe Trp Glu Ser Val
 740 745 750
 Phe Thr Gly Leu Thr His Ile Asp Ala His Phe Leu Ser Gln Thr Lys
 755 760 765
 Gln Ala Gly Asp Asn Phe Pro Tyr Leu Val Ala Tyr Gln Ala Thr Val
 770 775 780
 Cys Ala Arg Ala Gln Ala Pro Pro Pro Ser Trp Asp Gln Met Trp Lys
 785 790 795 800
 Cys Leu Ile Arg Leu Lys Pro Thr Leu His Gly Pro Thr Pro Leu Leu
 805 810 815
 Tyr Arg Leu Gly Ala Val Gln Asn Glu Val Thr Leu Thr His Pro Ile
 820 825 830

-42-

Thr Lys Tyr Ile Met Ala Cys Met Ser Ala Asp Leu Glu Val Val Thr
 835 840 845
 Ser Thr Trp Val Leu Val Gly Gly Val Leu Ala Ala Leu Ala Ala Tyr
 850 855 860
 Cys Leu Thr Thr Gly Ser Val Val Ile Val Gly Arg Ile Ile Leu Ser
 865 870 875 880
 Gly Arg Pro Ala Ile Val Pro Asp Arg Glu Leu Leu Tyr Gln Glu Phe
 885 890 895
 Asp Glu Met Glu Glu Cys Ala Ser His Leu Pro Tyr Ile Glu Gln Gly
 900 905 910
 Met Gln Leu Ala Glu Gln Phe Lys Gln Lys Ala Leu Gly Leu Leu Gln
 915 920 925
 Thr Ala Thr Lys Glu Ala Glu Ala Ala Ala Pro Val Val Glu Ser Lys
 930 935 940
 Trp Arg Ala Leu Glu Thr Phe Trp Ala Lys His Met Trp Asn Phe Ile
 945 950 955 960
 Ser Gly Ile Gln Tyr Leu Ala Gly Leu Ser Thr Leu Pro Gly Asn Pro
 965 970 975
 Ala Ile Ala Ser Leu Met Ala Phe Thr Ala Ser Ile Thr Ser Pro Leu
 980 985 990
 Thr Thr Gln Ser Thr Leu Leu Phe Asn Ile Leu Gly Gly Trp Val Ala
 995 1000 1005
 Ala Gln Leu Ala Pro Pro Ser Ala Ala Ser Ala Phe Val Gly Ala Gly
 1010 1015 1020
 Ile Ala Gly Ala Ala Val Gly Ser Ile Gly Leu Gly Lys Val Leu Val
 1025 1030 1035 1040
 Asp Ile Leu Ala Gly Tyr Gly Ala Gly Val Ala Gly Ala Leu Val Ala
 1045 1050 1055
 Phe Lys Val Met Ser Gly Glu Met Pro Ser Thr Glu Asp Leu Val Asn
 1060 1065 1070
 Leu Leu Pro Ala Ile Leu Ser Pro Gly Ala Leu Val Val Gly Val Val
 1075 1080 1085
 Cys Ala Ala Ile Leu Arg Arg His Val Gly Pro Gly Glu Gly Ala Val
 1090 1095 1100
 Gln Trp Met Asn Arg Leu Ile Ala Phe Ala Ser Arg Gly Asn His Val
 1105 1110 1115 1120
 Ser Pro Thr His Tyr Val Pro Glu Ser Asp Ala Ala Ala Arg Val Thr
 1125 1130 1135
 Gln Ile Leu Ser Ser Leu Thr Ile Thr Gln Leu Leu Lys Arg Leu His
 1140 1145 1150

-43-

Gln Trp Ile Asn Glu Asp Cys Ser Thr Pro Cys Ser Gly Ser Trp Leu
 1155 1160 1165
 Arg Asp Val Trp Asp Trp Ile Cys Thr Val Leu Thr Asp Phe Lys Thr
 1170 1175 1180
 Trp Leu Gln Ser Lys Leu Leu Pro Gln Leu Pro Gly Val Pro Phe Phe
 1185 1190 1195 1200
 Ser Cys Gln Arg Gly Tyr Lys Gly Val Trp Arg Gly Asp Gly Ile Met
 1205 1210 1215
 Gln Thr Thr Cys Pro Cys Gly Ala Gln Ile Thr Gly His Val Lys Asn
 1220 1225 1230
 Gly Ser Met Arg Ile Val Gly Pro Lys Thr Cys Ser Asn Thr Trp His
 1235 1240 1245
 Gly Thr Phe Pro Ile Asn Ala Tyr Thr Gly Pro Cys Thr Pro Ser
 1250 1255 1260
 Pro Ala Pro Asn Tyr Ser Arg Ala Leu Trp Arg Val Ala Ala Glu Glu
 1265 1270 1275 1280
 Tyr Val Glu Val Thr Arg Val Gly Asp Phe His Tyr Val Thr Gly Met
 1285 1290 1295
 Thr Thr Asp Asn Val Lys Cys Pro Cys Gln Val Pro Ala Pro Glu Phe
 1300 1305 1310
 Phe Ser Glu Val Asp Gly Val Arg Leu His Arg Tyr Ala Pro Ala Cys
 1315 1320 1325
 Arg Pro Leu Leu Arg Glu Glu Val Thr Phe Gln Val Gly Leu Asn Gln
 1330 1335 1340
 Tyr Leu Val Gly Ser Gln Leu Pro Cys Glu Pro Glu Pro Asp Val Ala
 1345 1350 1355 1360
 Val Leu Thr Ser Met Leu Thr Asp Pro Ser His Ile Thr Ala Glu Thr
 1365 1370 1375
 Ala Lys Arg Arg Leu Ala Arg Gly Ser Pro Pro Ser Leu Ala Ser Ser
 1380 1385 1390
 Ser Ala Ser Gln Leu Ser Ala Pro Ser Leu Lys Ala Thr Cys Thr Thr
 1395 1400 1405
 His His Val Ser Pro Asp Ala Asp Leu Ile Glu Ala Asn Leu Leu Trp
 1410 1415 1420
 Arg Gln Glu Met Gly Gly Asn Ile Thr Arg Val Glu Ser Glu Asn Lys
 1425 1430 1435 1440
 Val Val Val Leu Asp Ser Phe Asp Pro Leu Arg Ala Glu Glu Asp Glu
 1445 1450 1455
 Arg Glu Val Ser Val Pro Ala Glu Ile Leu Arg Lys Ser Lys Lys Phe
 1460 1465 1470

-44-

Pro Ala Ala Met Pro Ile Trp Ala Arg Pro Asp Tyr Asn Pro Pro Leu
 1475 1480 1485
 Leu Glu Ser Trp Lys Asp Pro Asp Tyr Val Pro Pro Val Val His Gly
 1490 1495 1500
 Cys Pro Leu Pro Pro Ile Lys Ala Pro Pro Ile Pro Pro Pro Arg Arg
 1505 1510 1515 1520
 Lys Arg Thr Val Val Leu Thr Glu Ser Ser Ser Val Ser Ala Leu Ala
 1525 1530 1535
 Glu Leu Ala Thr Lys Thr Phe Gly Ser Ser Glu Ser Ser Ala Val Asp
 1540 1545 1550
 Ser Gly Thr Ala Thr Ala Leu Pro Asp Gln Ala Ser Asp Asp Gly Asp
 1555 1560 1565
 Lys Gly Ser Asp Val Glu Ser Tyr Ser Ser Met Pro Pro Leu Glu Gly
 1570 1575 1580
 Glu Pro Gly Asp Pro Asp Leu Ser Asp Gly Ser Trp Ser Thr Val Ser
 1585 1590 1595 1600
 Glu Glu Ala Ser Glu Asp Val Val Cys Cys Ser Met Ser Tyr Thr Trp
 1605 1610 1615
 Thr Gly Ala Leu Ile Thr Pro Cys Ala Ala Glu Glu Ser Lys Leu Pro
 1620 1625 1630
 Ile Asn Ala Leu Ser Asn Ser Leu Leu Arg His His Asn Met Val Tyr
 1635 1640 1645
 Ala Thr Thr Ser Arg Ser Ala Gly Leu Arg Gln Lys Lys Val Thr Phe
 1650 1655 1660
 Asp Arg Leu Gln Val Leu Asp Asp His Tyr Arg Asp Val Leu Lys Glu
 1665 1670 1675 1680
 Met Lys Ala Lys Ala Ser Thr Val Lys Ala Lys Leu Leu Ser Val Glu
 1685 1690 1695
 Glu Ala Cys Lys Leu Thr Pro Pro His Ser Ala Lys Ser Lys Phe Gly
 1700 1705 1710
 Tyr Gly Ala Lys Asp Val Arg Asn Leu Ser Ser Lys Ala Val Asn His
 1715 1720 1725
 Ile His Ser Val Trp Lys Asp Leu Leu Glu Asp Thr Val Thr Pro Ile
 1730 1735 1740
 Asp Thr Thr Ile Met Ala Lys Asn Glu Val Phe Cys Val Gln Pro Glu
 1745 1750 1755 1760
 Lys Gly Gly Arg Lys Pro Ala Arg Leu Ile Val Phe Pro Asp Leu Gly
 1765 1770 1775
 Val Arg Val Cys Glu Lys Met Ala Leu Tyr Asp Val Val Ser Thr Leu
 1780 1785 1790

-45-

Pro Gln Val Val Met Gly Ser Ser Tyr Gly Phe Gln Tyr Ser Pro Gly
 1795 1800 1805
 Gln Arg Val Glu Phe Leu Val Asn Thr Trp Lys Ser Lys Lys Asn Pro
 1810 1815 1820
 Met Gly Phe Ser Tyr Asp Thr Arg Cys Phe Asp Ser Thr Val Thr Glu
 1825 1830 1835 1840
 Asn Asp Ile Arg Val Glu Glu Ser Ile Tyr Gln Cys Cys Asp Leu Ala
 1845 1850 1855
 Pro Glu Ala Arg Gln Ala Ile Lys Ser Leu Thr Glu Arg Leu Tyr Ile
 1860 1865 1870
 Gly Gly Pro Leu Thr Asn Ser Lys Gly Gln Asn Cys Gly Tyr Arg Arg
 1875 1880 1885
 Cys Arg Ala Ser Gly Val Leu Thr Thr Ser Cys Gly Asn Thr Leu Thr
 1890 1895 1900
 Cys Tyr Leu Lys Ala Ser Ala Ala Cys Arg Ala Ala Lys Leu Gln Asp
 1905 1910 1915 1920
 Cys Thr Met Leu Val Asn Gly Asp Asp Leu Val Val Ile Cys Glu Ser
 1925 1930 1935
 Ala Gly Thr Gln Glu Asp Ala Ala Ser Leu Arg Val Phe Thr Glu Ala
 1940 1945 1950
 Met Thr Arg Tyr Ser Ala Pro Pro Gly Asp Pro Pro Gln Pro Glu Tyr
 1955 1960 1965
 Asp Leu Glu Leu Ile Thr Ser Cys Ser Ser Asn Val Ser Val Ala His
 1970 1975 1980
 Asp Ala Ser Gly Lys Arg Val Tyr Tyr Leu Thr Arg Asp Pro Thr Thr
 1985 1990 1995 2000
 Pro Leu Ala Arg Ala Ala Trp Glu Thr Ala Arg His Thr Pro Val Asn
 2005 2010 2015
 Ser Trp Leu Gly Asn Ile Ile Met Tyr Ala Pro Thr Leu Trp Ala Arg
 2020 2025 2030
 Met Ile Leu Met Thr His Phe Phe Ser Ile Leu Leu Ala Gln Glu Gln
 2035 2040 2045
 Leu Glu Lys Ala Leu Asp Cys Gln Ile Tyr Gly Ala Cys Tyr Ser Ile
 2050 2055 2060
 Glu Pro Leu Asp Leu Pro Gln Ile Ile Glu Arg Leu His Gly Leu Ser
 2065 2070 2075 2080
 Ala Phe Ser Leu His Ser Tyr Ser Pro Gly Glu Ile Asn Arg Val Ala
 2085 2090 2095
 Ser Cys Leu Arg Lys Leu Gly Val Pro Pro Leu Arg Val Trp Arg His
 2100 2105 2110

-46-

Arg Ala Arg Ser Val Arg Ala Arg Leu Leu Ser	Gln Gly Gly Arg Ala		
2115	2120	2125	
Ala Thr Cys Gly Lys Tyr Leu Phe Asn Trp Ala Val Lys Thr Lys Leu			
2130	2135	2140	
Lys Leu Thr Pro Ile Pro Ala Ala Ser Arg	Leu Asp Leu Ser Gly Trp		
2145	2150	2155	2160
Phe Val Ala Gly Tyr Ser Gly Gly Asp Ile Tyr His Ser Leu Ser Arg			
2165	2170	2175	
Ala Arg Pro Arg Trp Phe Met Leu Cys Leu	Leu Leu Leu Ser Val Gly		
2180	2185	2190	
Val Gly Ile Tyr Leu Leu Pro Asn Arg			
2195	2200		

(3) INFORMAZIONI PER SEQ ID NO: 3

(i) CARATTERISTICHE DELLA SEQUENZA:

- (A) LUNGHEZZA: 26 nucleotidi
- (B) TIPO: acido nucleico
- (C) NUMERO DI CATENE: singola
- (D) CONFIGURAZIONE: lineare
- (ii) TIPO DI MOLECOLA: DNA sintetico
- (iii) IPOTETICA: No
- (iv) ANTISENSO: No
- (vii) FONTE IMMEDIATA: sintetizzatore di oligonucleotidi
- (ix) CARATTERISTICHE:
 - (A) NOME: oligo a
 - (C) METODO DI IDENTIFICAZIONE: gel di poliacrilammide
- (xi) DESCRIZIONE DELLA SEQUENZA: SEQ ID NO: 3

GCCGAGATGC CATCTTCAAA CAGTTC

26

(4) INFORMAZIONI SULLA SEQUENZA: SEQ ID NO: 4

(i) CARATTERISTICHE DELLA SEQUENZA:

- (A) LUNGHEZZA: 24 nucleotidi
- (B) TIPO: acido nucleico
- (C) NUMERO DI CATENE: singola
- (D) CONFIGURAZIONE: lineare
- (ii) TIPO DI MOLECOLA: DNA sintetico
- (iii) IPOTETICA: No
- (iv) ANTISENSO: No
- (vii) FONTE IMMEDIATA: sintetizzatore di oligonucleotidi
- (ix) CARATTERISTICHE:

(3) INFORMATION FOR SEQ ID NO: 3:

(i) SEQUENCE CHARACTERISTICS

(A) LENGTH: 26 nucleotides
(B) TYPE: nucleic acid
(C) NUMBER OF CHAINS: single
(D) CONFIGURATION: linear

(ii) MOLECULE TYPE: synthetic DNA

(iii) HYPOTHETICAL: No

(iv) ANTISENSE: No

(vii) IMMEDIATE SOURCE: oligonucleotide synthesizer

(ix) CHARACTERISTICS:

(A) NAME: oligo a
(C) IDENTIFICATION METHOD: Polyacrylamide gel

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3

GCCGAGATGC CATCTTCAAA CAGTTC

26

(4) INFORMATION FOR SEQ ID NO: 4:

(i) SEQUENCE CHARACTERISTICS

(A) LENGTH: 24 nucleotides
(B) TYPE: nucleic acid
(C) NUMBER OF CHAINS: single
(D) CONFIGURATION: linear

(ii) MOLECULE TYPE: synthetic DNA

(iii) HYPOTHETICAL: No

(iv) ANTISENSE: No

(vii) IMMEDIATE SOURCE: oligonucleotide synthesizer

(ix) CHARACTERISTICS:

(A) NAME: oligo b
(C) IDENTIFICATION METHOD: Polyacrylamide gel

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4

GTGTACAACA AGGTCCATAT CACC

24

(5) INFORMATION FOR SEQ ID NO: 5:

(i) SEQUENCE CHARACTERISTICS
(A) LENGTH: 24 nucleotides
(B) TYPE: nucleic acid
(C) NUMBER OF CHAINS: single
(D) CHARACTERISTICS: linear

(ii) MOLECULE TYPE: synthetic DNA

(iii) HYPOTHETICAL: No

(iv) ANTISENSE: No

(vii) IMMEDIATE SOURCE: oligonucleotide synthesizer

(ix) CHARACTERISTICS:
(A) NAME: oligo c
(C) IDENTIFICATION METHOD: Polyacrylamide gel

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5

GGTCTTTCTG AACGGGATAT AAAC

24

(6) INFORMATION FOR SEQ ID NO: 6:

(i) SEQUENCE CHARACTERISTICS
(A) LENGTH: 31 nucleotides
(B) TYPE: nucleic acid
(C) NUMBER OF CHAINS: single
(D) CONFIGURATION: linear

(ii) MOLECULE TYPE: synthetic DNA

(iii) HYPOTHETICAL: No

(iv) ANTISENSE: No

(vii) IMMEDIATE SOURCE: oligonucleotide synthesizer

(ix) CHARACTERISTICS:
(A) NAME: 5'-5B
(C) IDENTIFICATION METHOD: Polyacrylamide gel

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6

AAGGATCCAT GTCAATGTCC TACACATGGA C

31

(7) INFORMATION FOR SEQ ID NO: 7:

(i) SEQUENCE CHARACTERISTICS

(A) LENGTH: 36 nucleotides
(B) TYPE: nucleic acid
(C) NUMBER OF CHAINS: single
(D) CONFIGURATION: linear
(ii) MOLECULE TYPE: synthetic DNA
(iii) HYPOTHETICAL: No
(iv) ANTISENSE: No
(vii) IMMEDIATE SOURCE: oligonucleotide synthesizer
(ix) CHARACTERISTICS:
(A) NAME: 3'-5B
(C) IDENTIFICATION METHOD: Polyacrylamide gel
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7

AATATTCGAA TTCATCGGTT GGGGAGCAGG TAGATG

36

(8) INFORMATION FOR SEQ ID NO: 8:
(i) SEQUENCE CHARACTERISTICS
(A) LENGTH: 22 nucleotides
(B) TYPE: nucleic acid
(C) NUMBER OF CHAINS: single
(D) CHARACTERISTICS: linear
(ii) MOLECULE TYPE: synthetic DNA
(iii) HYPOTHETICAL: No
(iv) ANTISENSE: No
(vii) IMMEDIATE SOURCE: oligonucleotide synthesizer
(ix) CHARACTERISTICS:
(A) NAME: Dpr1
(C) IDENTIFICATION METHOD: Polyacrylamide gel
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8

TGGCTGGCAA GGCACACAGG CT

22

(9) INFORMATION FOR SEQ ID NO: 9:
(i) SEQUENCE CHARACTERISTICS
(A) LENGTH: 20 nucleotides

(B) TYPE: nucleic acid
(C) NUMBER OF CHAINS: single
(D) CONFIGURATION: linear
(ii) MOLECULE TYPE: synthetic DNA
(iii) HYPOTHETICAL: No
(iv) ANTISENSE: No
(vii) IMMEDIATE SOURCE: oligonucleotide synthesizer
(ix) CONFIGURATION:
(A) NAME: Dpr2
(C) IDENTIFICATION METHOD: Polyacrylamide gel
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9

AGGCAGGGTA GATCTATGTC

20

(10) INFORMATION FOR SEQ ID NO: 10:
(i) SEQUENCE CHARACTERISTICS
(A) LENGTH: 20 nucleotides
(B) TYPE: nucleic acid
(C) NUMBER OF CHAINS: single
(D) CONFIGURATION: linear
(ii) MOLECULE TYPE: synthetic DNA
(iii) HYPOTHETICAL: No
(iv) ANTISENSE: No
(vii) IMMEDIATE SOURCE: oligonucleotide synthesizer
(ix) CHARACTERISTICS:
(A) NAME: NS5B-5' (1)
(C) IDENTIFICATION METHOD: Polyacrylamide gel
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10

TCAATGTCCT ACACATGGAC

20

(11) INFORMATION FOR SEQ ID NO: 11:
(i) SEQUENCE CHARACTERISTICS
(A) LENGTH: 38 nucleotides
(B) TYPE: nucleic acid

- (C) NUMBER OF CHAINS: single
- (D) CONFIGURATION: linear
- (ii) MOLECULE TYPE: synthetic DNA
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (vii) IMMEDIATE SOURCE: oligonucleotide synthesizer
- (ix) CHARACTERISTICS:
 - (A) NAME: HCVA-13
 - (C) IDENTIFICATION METHOD: Polyacrylamide gel
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11

GATCTCTAGA TCATCGGTTG GGGGAGGAGG TAGATGCC 38

(12) INFORMATION FOR SEQ ID NO: 12:

- (i) SEQUENCE CHARACTERISTICS
 - (A) LENGTH: 399 nucleotides
 - (B) TYPE: nucleic acid
 - (C) NUMBER OF CHAINS: single
 - (D) CONFIGURATION: linear
- (ii) MOLECULE TYPE: mRNA
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Rattus Norvegicus
 - (C) STRAIN: Sprague-Dawley
- (vii) IMMEDIATE SOURCE: pT7-7 (DCoH)
- (ix) CHARACTERISTICS:
 - (A) NAME: D-RNA
 - (C) IDENTIFICATION METHOD: Polyacrylamide gel
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12

CLAIMS

1. Method for reproducing *in vitro* the RNA-dependent RNA polymerase activity or the terminal nucleotidyl transferase activity encoded by hepatitis C virus (HCV), wherein sequences containing NS5B (SEQ ID NO: 1) are used in the reaction mixture.

2. Method for reproducing *in vitro* the RNA-dependent RNA polymerase activity encoded by HCV according to Claim 1, wherein NS5B is incorporated in the reaction mixture as NS2-NS5B precursor, said precursor generating, by means of multiple proteolytic events that occur in the overproducing organism, an enzymatically active form of NS5B.

3. Method for reproducing *in vitro* the terminal nucleotidyl transferase activity coded for HCV according to Claim 1, wherein NS5B is incorporated in the reaction mixture as NS2-NS5B precursor, said precursor generating, by means of multiple proteolytic events that occur in the overproducing organism, an enzymatically active form of NS5B.

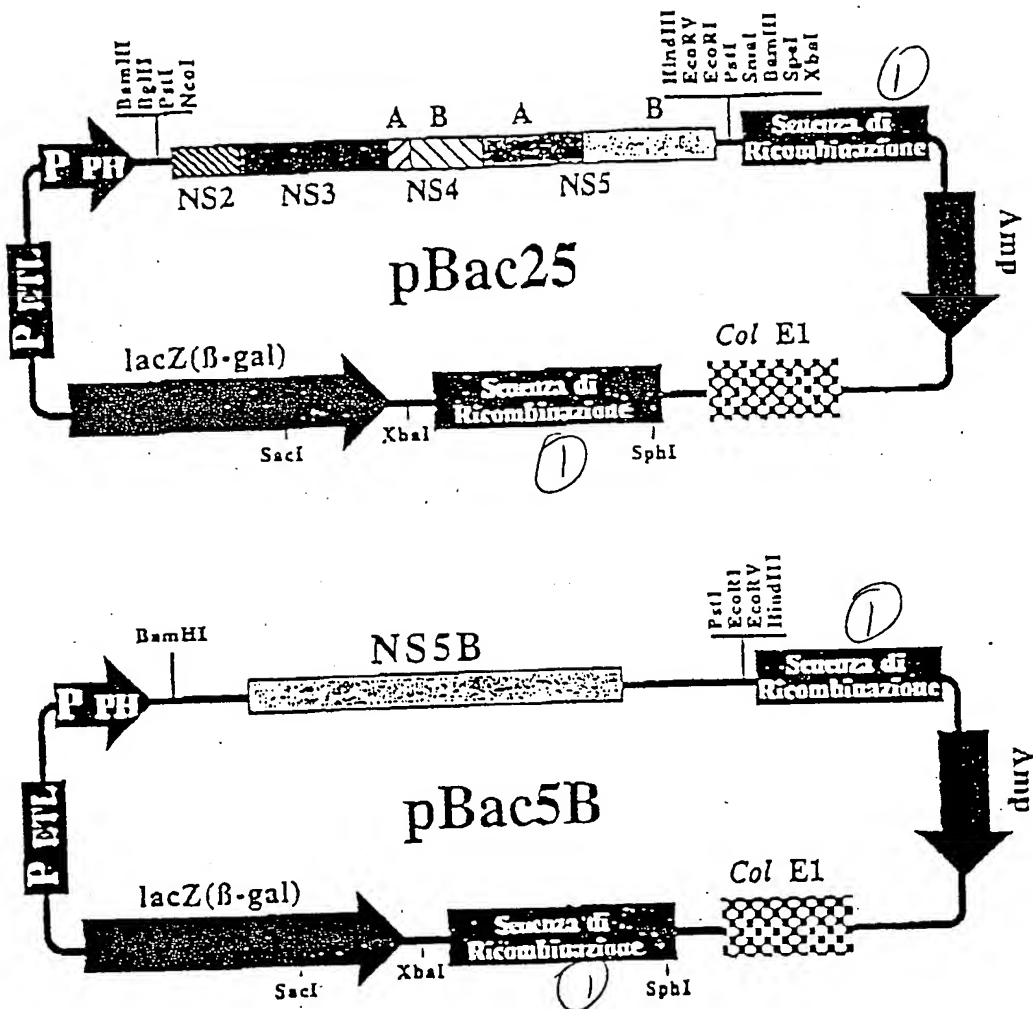
4. Composition of matter containing NS5B sequences according to Claims 1 through 3.

5. Composition of matter according to Claim 4, comprising the proteins whose sequences are described in SEQ ID NO: 1, in sequences contained therein or derived therefrom.

6. Use of the compositions of matter according to Claims 4 and 5 to set up an enzymatic test capable of selecting, for therapeutic purposes, compounds that inhibit the enzymatic activity associated with NS5B.

7. Method for reproducing *in vitro* the RNA-dependent RNA polymerase and terminal nucleotidyl transferase activities, compositions of matter and use of said compositions of matter to set up an enzymatic test capable of selecting, for therapeutic purposes, compounds that inhibit the enzymatic activities associated with NS5B, according to the above description, examples and claims.

ISTITUTO DI RICERCHE DI BIOLOGIA MOLECOLARE ANGELETTI S.p.A.



[Key]

1. Recombination sequence

P_{ETL} = Promoter of the gene coding for PCNA protein

P_{PH} = Promoter of the polyhedrin gene

Amp = Gene coding for the β-lactamase enzyme
(ampicillin resistance)

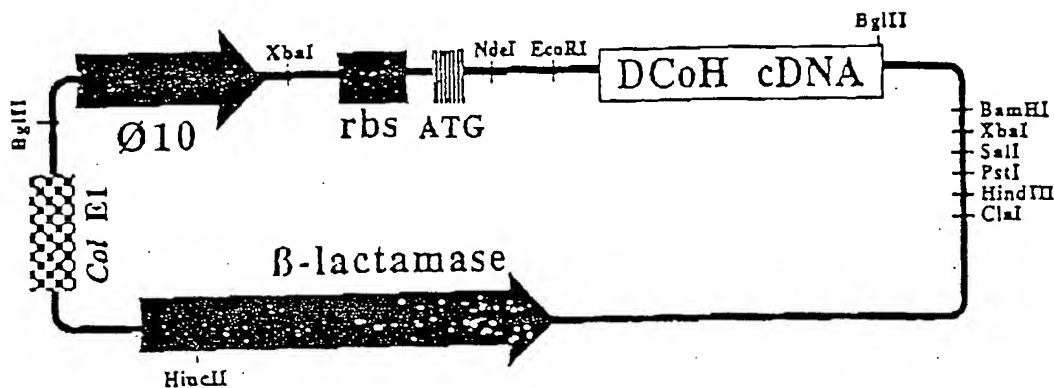
LacZ (β-gal) = Gene coding for the β-galactosidase enzyme

Col E1 = pBR322 replication origin

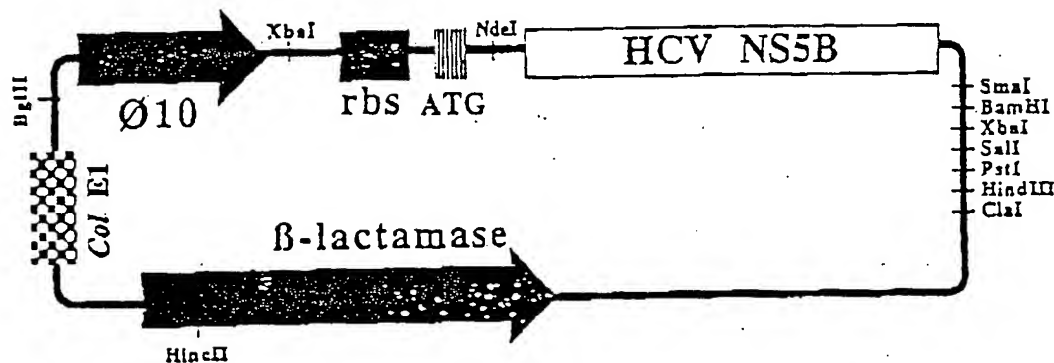
Figure 1

PT7-7 (DCoH)

pT7-7(DCoH)



pT7-7(NS5B)



ø10 = Bacteriophage T7 ø10 promoter
rbs = Shine-Dalgarno ribosome binding site
ATG = Translation initiation site of the protein coded by the bacteriophage T7 gene 10
β-lactamase = gene coding for the β-lactamase enzyme (ampicillin resistance)
Col E1 = pBR322 replication origin

Figure 2

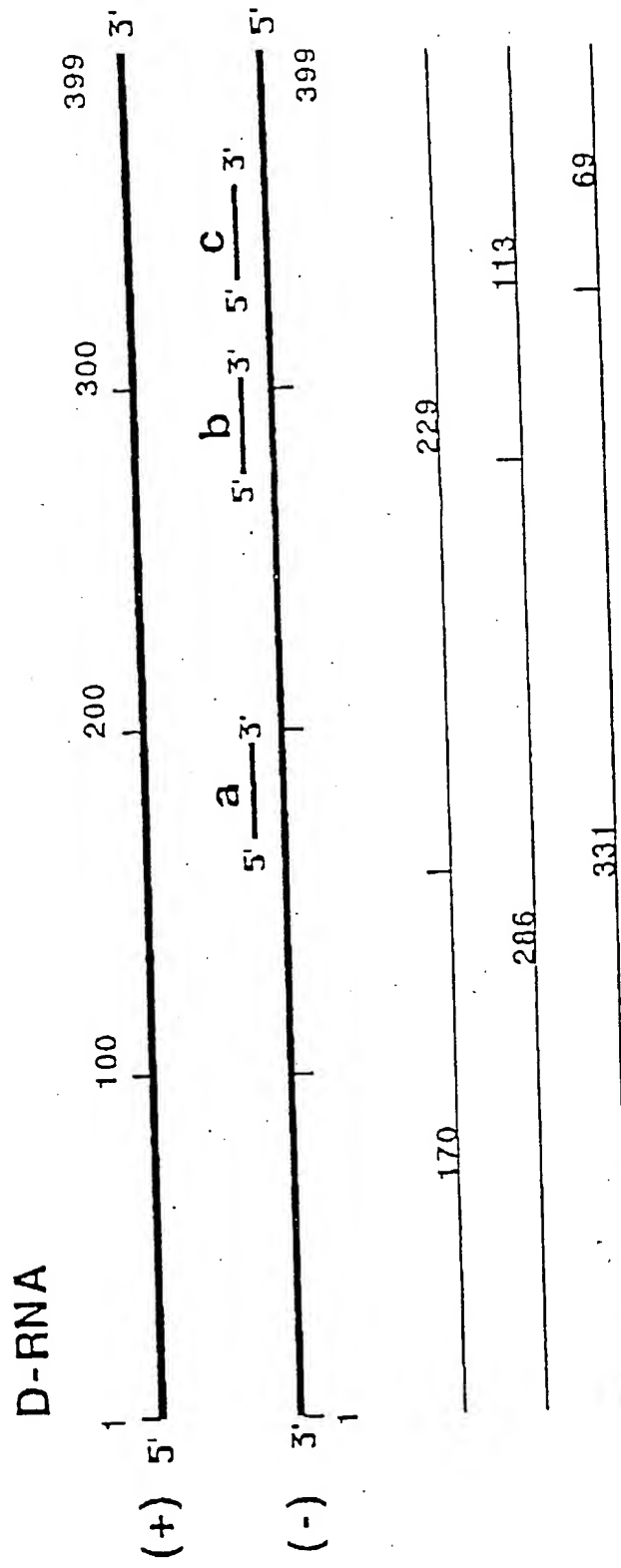


FIG. 3

MINISTRY OF INDUSTRY, COMMERCE AND ARTISANRY
GENERAL ADMINISTRATION OF PRODUCTIVE DEVELOPMENT AND
COMPETITIVENESS
ITALIAN PATENT AND TRADEMARK OFFICE

PATENT FOR INDUSTRIAL INVENTION

No. 01278077

This patent was granted for the invention that was the subject of the application below:

Applica- tion No.	Year	U.P.I.C.A.	Date of Presenta- tion of Applica- tion	Classifi- cation
000343	95	ROME	5/25/1995	C12Q

HOLDER: ISTITUTO DI RICERCHE DI BIOLOGIA
MOLECOLARE S.p.A.
IN POMEZIA (ROME)

AGENT: DE BENEDETTI, FABRIZIO

ADDRESS: SOCIETÀ ITALIANA BREVETTI SPA
P.ZA DI PIETRA 39
00100 ROME

TITLE: METHOD FOR REPRODUCING IN VITRO THE RNA-
DEPENDENT RNA POLYMERASE AND TERMINAL
NUCLEOTIDYL TRANSFERASE ACTIVITIES
ENCODED BY HEPATITIS C VIRUS (HCV)

INVENTOR: DE FRANCESCO, RAFFAELE
BEHRENS, SVEN ERIK
TOMEI LICIA

[seal]
Ministry of Industry,
Commerce and Artisanry
Italian Patent and
Trademark Office
[canceled tax stamp]

Rome, NOVEMBER 17, 1997

DIRECTOR OF DIVISION V

Signed: GIOVANNA MORELLI

CERTIFIED TRUE COPY OF THE ORIGINAL
[Illegible signature]

Issued on December 3, 1997
The Director of the UPICA
[Illegible signature]